

1/5

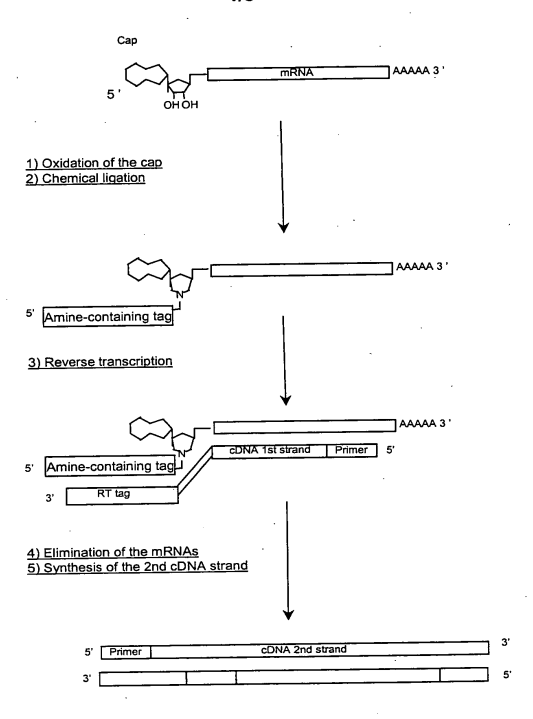


Figure 1

APPROVED O. FIG.
BY QL/38 SUFFCLASS

DEMETSMAN

Minimum signal peptide score	false positive rate	false negative rate	proba(0.1)	proba(0.2)
3.5	0.121	0.036	0.467	0.664
4	0.096	0.06	0.519	0.708
4.5	0.078	0.079	0.565	0.745
5	0.062	0.098	0.615	0.782
5.5	0.05	0.127	0.659	0.813
6	0.04	0.163	0.694	0.836
6.5	0.033	0.202	0.725	0.855
7	0.025	0.248	0.763	0.878
7.5	0.021	0.304	0.78	0.889
8	0.015	0.368	0.816	0.909
8.5	0.012	0.418	0.836	0.92
9	0.009	0.512	0.856	0.93
9.5	0.007	0.581	0.863	0.934
10	0.006	0.679	0.835	0.919

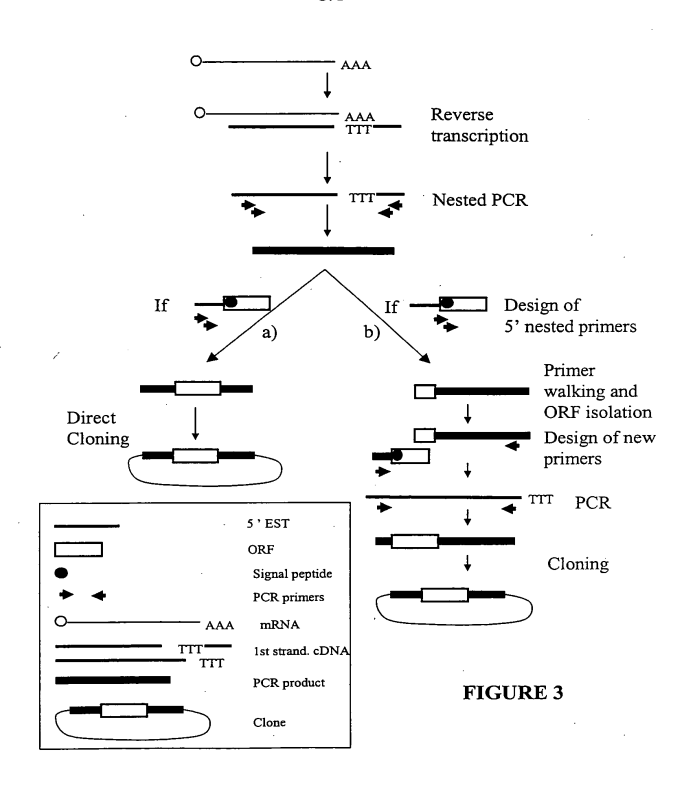
Figure 2

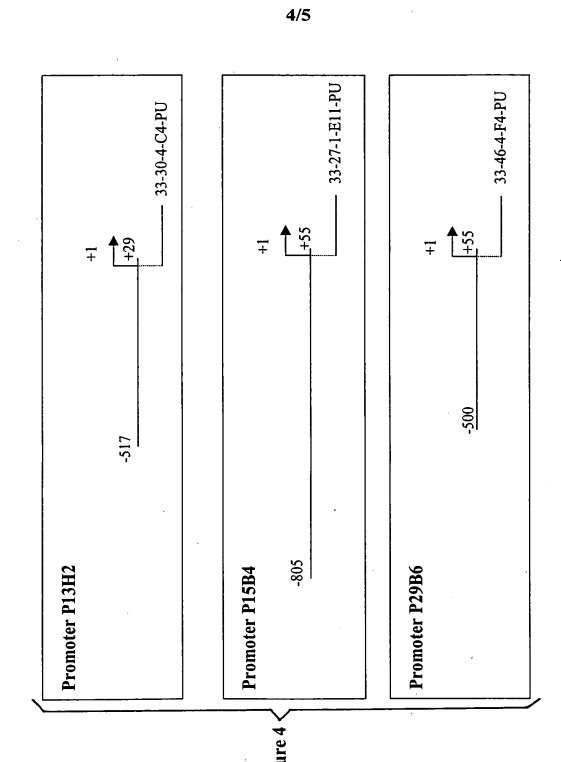
APPROVED C

CHAFTSMAN

FIR

CLASS





AFROVED C. FIG.

BY DHAFTSMAN

	Jealon characteristic	2		nnaiac		SIICS
Step	Program	Strand	Parameters	Identity (%)	Length (bp)	Comments
miscellanaeous	FASTA	both		06	15	
tRNA	FASTA	both		80	09	
rRNA	BLASTN	both	S=108	80	40	
mtRNA	BLASTN	both	S=108	- 80	40	
Procaryotic	BLASTN	both	S=144	06	40	
Fungal	BLASTN	both	S=144	06	40	
Alu	BLASTN	both	S=72, B=5	20	40	max 5 matches, masking
L1	BLASTN	both	S=72, B=5	20	40	max 5 matches, masking
Repeats	BLASTN	both	S=72	02	40	masking
Aviod	DI ACTON	Ç	W=6, S=10,	8	,	
Loigh	DLAS I ZN	do ₁	E=1000, N=-12	90	10	in the last 100 nucleotides
Polyadenylation signal	-	top	AATAAA al	AATAAA allowing 1 mismatch	atch	in the 50 nucleotides preceding the 5' end of the polA
	BLASTN then					first BLASTN and then FASTA
Vertebrate	FASTA	poth	•	90 then 70	30	on maching sequences
ESTS	BLAST2N	both	•	06	30	
Genesed	BLASTN	both	W=8, B=10	06	30	
ORF	BLASTP	top	W=8, B=10	ı	1	on ORF proteins, max 10 matches
Proteins	BLASTX	top	E = 0.001	70	30	

Parameters used for each step of cDNA analysis